



REVIEW

Vestibular animal models: contributions to understanding physiology and disease

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Abstract Our knowledge of the vestibular sensory system, its functional significance for gaze and posture stabilization, and its capability to ensure accurate spatial orientation perception and spatial navigation has greatly benefitted from experimental approaches using a variety of vertebrate species. This review summarizes the attempts to establish the roles of semicircular canal and otolith endorgans in these functions followed by an overview of the most relevant fields of vestibular research including major findings that have advanced our understanding of how this system exerts its influence on reflexive and cognitive challenges encountered during daily life. In particular, we highlight the contributions of different animal models and the advantage of using a comparative research approach. Cross-species comparisons have established that the morpho-physiological properties underlying vestibular signal processing are evolutionarily inherent, thereby disclosing general principles. Based on the documented success of this approach, we suggest that

future research employing a balanced spectrum of standard animal models such as fish/frog, mouse and primate will optimize our progress in understanding vestibular processing in health and disease. Moreover, we propose that this should be further supplemented by research employing more “exotic” species that offer unique experimental access and/or have specific vestibular adaptations due to unusual locomotor capabilities or lifestyles. Taken together this strategy will expedite our understanding of the basic principles underlying vestibular computations to reveal relevant translational aspects. Accordingly, studies employing animal models are indispensable and even mandatory for the development of new treatments, medication and technical aids (implants) for patients with vestibular pathologies.

Keywords Otolith organ · Semicircular canal · Sensory-motor processing · Motion perception · Gaze stabilization · Vestibulo-ocular reflex

This manuscript is part of a supplement sponsored by the German Federal Ministry of Education and Research within the funding initiative for integrated research and treatment centers.

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Introduction

Scientific research on the vestibular system has benefitted from studies on a wide variety of vertebrate species [1]. Systematic investigations of inner ear endorgans in fishes, amphibians and birds, performed over a century ago, revealed many details of the structure/function of this sensory system and established that it is remarkably preserved across vertebrate phylogeny [2, 3]. This organizational conservation emphasizes that the ability to detect and encode body motion for gaze and posture stabilization as well as for orientation and navigation in space is essential across vertebrates [4, 5]. In addition, the almost identical structure of sensory endorgans, neuronal pathways and

central circuits in different vertebrates provides the ability to make general conclusions regarding the signaling properties and computational capabilities of the neuronal components of vestibular pathways. Moreover, species-specific features, related to different locomotor strategies/dynamics, particular lifestyles or eco-physiological habitats, offer the opportunity to evaluate the capacity of the system to adapt to new challenges. Thus, understanding the particularities and similarities of vestibular signal processing in different vertebrates have provided an essential opportunity to identify conceptual principles that coincide with the behavioral repertoire and performance. This knowledge has in turn facilitated our understanding of the mechanistic operations required for stabilizing gaze and posture and yielded insight into substrates and processes underlying different pathologies and potential treatments.

Notably, based on their evolutionary proximity to humans, non-human primates have become a standard model for furthering our knowledge of basic vestibular processing and advancing translational research. In addition, advances in the generation of mouse lines with defined genetic backgrounds and/or transgenic lines, such as CRE, combined with viral-based optogenetics have provided new opportunities to probe the functional circuitry of vestibular pathways and gain insight into vestibular diseases or age-related impairments [6–8]. However, it is also important to emphasize that work across a wider variety of different vertebrate models has improved our understanding of vestibular processing. Research encompassing a combination of standard models and less widely used “exotic animals” with particular motion repertoires and/or unique experimental advantages have proven advantageous. This review highlights recent progress that has been made toward understanding the fundamental physiological principles of vestibular processing using different animal models as well as how cellular and circuit properties are altered under pathophysiological conditions.

Animal models and major topics in vestibular research

Historical overview

The vestibular endorgans are located within the petrosal bone in close vicinity to the cochlea. This hidden location proximal to the auditory organ and the difficult access considerably delayed our understanding of the functional role of these inner ear organs. It was only in 1824, that Flourens [9] conducted the first systematic behavioral studies on the vestibular system. He discovered that interrupting specific semicircular canals in different

vertebrate species, including pigeons and rabbits, produced direction-specific impairments of the equilibrium, walking and head movements. Interestingly, he speculated that the deficits were due to changes in hearing sensitivity, since the semicircular canals were then generally considered as part of the auditory organ. In fact, the vestibular system was only postulated to be a distinct sense independent of hearing in the late 19th century following systematic lesion experiments on frog, pigeon [2] and dog [10]. Both, Goltz [2] and Bechterew [10] concluded that the semicircular canals were distinct organs responsible for posture and equilibrium in three spatial orientations. This view was subsequently confirmed by studies in numerous other species including salamanders, pigeons, cockatoos and rabbits [3]. Concurrently, more theoretically based investigations determined that the semicircular canals sense head rotations [11–13].

As for the semicircular canals, the initially assumed auditory role of otolith organs also remained unchallenged for a long time. It is noteworthy that this historical progression in our understanding is reflected in existing terminology, since otolith literally means “hearing stone”. Theoretically based investigations had deduced that the otolith endorgans sense linear accelerations including head tilts [12, 13], yet experimental work in fish found that removal of the large, solid crystals (otoconia) covering the otolith sensory epithelia [14] significantly impaired their underwater hearing [15]. Experiments in terrestrial vertebrates were required to definitively establish that the otolith organs are responsible for ensuring stable posture and equilibrium in land-based animals [3, 16]. This conclusion was then furthered by experiments in fish indicating that, in this specific group of vertebrates, the saccule/lagena may also serve as a hearing organ [17], thus exerting a dual functional role.

In summary, a comparative approach including studies in vertebrate species from fish to mammals was essential to the progress that was made in the early vestibular research of the 19th century. In particular, the knowledge obtained following experimental lesions of semicircular canals, otoliths, and/or their nerves using different animal models has proven crucial for providing the important insight into species-specific adaptations of the endorgans and variations in their function relative to lifestyle, eco-physiological niche or locomotor dynamics.

Principles of mechanotransduction and hair cell dynamics

Head motion relative to space is detected and decomposed into individual vector components by semicircular canal and otolith organs [18]. The semicircular canals and otolith organs sense rotation, and linear motion or changes in head

position relative to the Earth's gravitation vector, respectively. The fluid-filled ducts of the semicircular canals enable the detection of angular acceleration by means of the fluid's inertia relative to sensory epithelia. The mechanistic principle of the otolith organs is based on the inertia of an otolithic structure covering the sensory epithelia [19]. While the spatial arrangement of the bilateral semicircular canals is largely conserved across vertebrates [20], comparative studies in fish, frogs and birds demonstrate that otolith organs can serve an auditory as well as vestibular function as a result of differences in the morpho-physiological properties of hair cells at a particular region of the otolithic epithelium [21]. Thus, in species such as frogs, fish and likely also in mammals, the otolith organs detect changes of the body position relative to the gravity vector as well as substrate/water vibrations [22]. Furthermore, the lagena, an otolith organ that is present in all non-mammalian vertebrates and monotremes [23], likely contributes to the sense of magnetoreception that allows birds to make use of the geomagnetic field for orientation and navigation [24].

Current evidence suggests that vestibular endorgans across species have adapted to changes in the environment and body mass [25], indicating that vestibular processing is influenced by the statistics of natural stimuli encountered in the sensory environment [26]. For example, the transition from water to land-based life has resulted in major changes in the natural sensory environment since the resistive hydrodynamic forces of an aquatic environment effectively dampen self-motion [27]. In addition, the longer and more flexible necks of amniotes can cause faster head movements [28]. Together, these factors suggest that terrestrial amniotes generally experience stimulation at higher amplitudes compared to mostly neckless fish or amphibian species.

During ontogeny, vestibular reflexes are generally present after hatching/birth in precocial animals such as larval fish, amphibian tadpoles or certain avian species or as soon as the respective neural circuitry and/or cellular properties are mature in altricial species [29]. While the developmental onset of otolith function is independent of animal size, semicircular canal functionality critically depends on tube dimensions as predicted by theoretical considerations [30] and verified by experimental data [31–34]. In particular, studies on the small larvae of fish and amphibians established the time course of onset, progression and maturation of motion-evoked semicircular canal-dependent reflexes after embryogenesis [34, 35]. This illustrates a general size dependency of semicircular canal reflexes and an important role for the spatial tuning of otolith-derived extraocular motor responses [36].

Hair cells in the vestibular system have long been a subject of functional studies using various vertebrate

species including bullfrog [37, 38], turtle, chinchilla and monkey (reviewed in [39]). While two types of hair cells (type I and II) are present in the inner ear of amniote vertebrates, anamniotes such as fish and amphibians possess only type II hair cells, which are exclusively contacted by bouton-like afferent terminals [39]. The evolutionary appearance of hair cells with a calyx-like afferent synapse (type I hair cells) in amniotes corresponds to a terrestrial lifestyle as well as with the appearance of flexible necks, since as noted above both factors likely contribute to higher frequency and acceleration head movements. This view concurs with the fact that responses of type I hair cells are considerably more dynamic than those of type II hair cells [39]. However, despite the absence of type I hair cells, frog otolith hair cells exhibit a similarly broad spectrum of response dynamics, suggesting that a calyx-like structure is no prerequisite for encoding high dynamic motion stimuli [40]. Recent studies in chinchilla and rodents [41] suggest that more detailed investigations of the complex type I hair cell calyx are required to fully understand its physiological implications/advantages.

Neural encoding strategies in the peripheral vestibular system

Previous studies have shown that afferents supplied by type I hair cells, which are found in amniotes, tend to be more irregular (Fig. 1a) in their resting discharges than afferents that contact type II hair cells [39]. This obvious dichotomy is compatible with the differential composition of ion conductances in vestibular ganglion cells (reviewed in [27]). Irregular afferents with calyx-like terminals have higher sensitivities and are better suited for the processing of natural motion stimuli in comparison to regular afferents (Fig. 1b) [26]. However, while there are clear physiological differences between irregular and regular afferent fibers in amniotes, it is notable that afferent response dynamics also vary in anamniotes. For example, a subclass of semicircular canal afferents in toadfish encodes angular acceleration [42], even though anamniotes have only type II hair cells.

Over the range of frequencies typically experienced during everyday behaviors (i.e., up to 20 Hz), semicircular canal afferents encode head velocity while otolith afferents encode linear acceleration in mammals ranging from mice to primates [43–45]. Quantification of afferent responses in primates has revealed important differences not only in the dynamics of regular versus irregular afferent activity (i.e., irregular afferents have higher sensitivities and phase leads as shown in Fig. 1b), but also in their information coding (Fig. 1c). In monkeys, regular canal afferents transmit twofold more information and are twice as sensitive for detecting head motion compared to irregular afferents [46]. Thus, regular and irregular canal afferents essentially

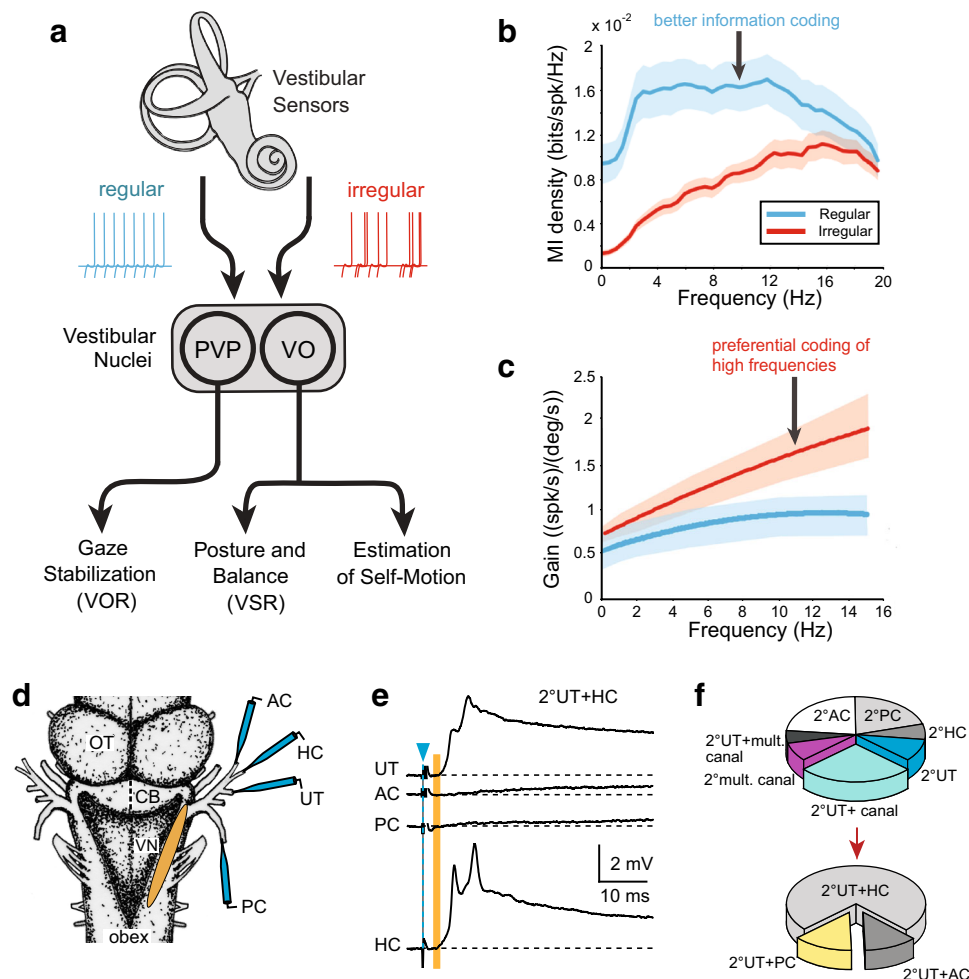


Fig. 1 Vestibular sensory-motor signal processing. **a** Schematic illustrating the two channels of input from labyrinthine nerve afferents (regular, irregular) and main subclasses of central vestibular neurons (PVP, VO) underlying vestibular reflexes (VOR, VSR) and self-motion perception. **b, c** Gain and mutual information density for regular and irregular vestibular afferent fibers. Population-averaged mutual information density curves (\pm SEM, **b**) and gains (\pm SEM, **c**) during random head rotations as function of frequency. **d–f**, Convergence of monosynaptic semicircular canal and otolith signals in frog 2°VNs; **d** schematic of an isolated frog whole brain depicting the electrical stimulation of individual labyrinthine nerve branches

and central vestibular recording area (orange); **e** 2°VNs, identified by monosynaptic (vertical orange bar) EPSPs (UT + HC) following separate stimulation (blue arrowhead) of the AC, HC, PC and UT nerve branches; **f** convergence pattern of utricular and semicircular canal nerve afferent inputs in identified 2°VNs. AC, PC, HC anterior, posterior vertical, horizontal semicircular canal, AP, BP amphibian, basal papilla, CB cerebellum, LA lagena, OT optic tectum, PVP position-vestibular-pause neuron, UT utricle, VN vestibular nuclei, VO vestibular-only neuron, VOR vestibulo-ocular reflex, VSR vestibulo-spinal reflex. **b, c, e, f** are based on data from [46] and [70], respectively

comprise two parallel information channels, one which encodes high-frequency stimuli with higher gains (i.e., irregular afferents), the other which transmits information about the detailed time course of the stimulus over the behaviorally significant frequency range (i.e., regular afferents). Interestingly, a different coding strategy is used by otolith afferents. Irregular otolith afferents are far more sensitive than regular otolith afferents, so much so that their differences in sensitivities are effectively compensated by differences in variability [45]. As a result, neuronal thresholds are independent of both stimulus frequency and resting discharge regularity.

Our basic understanding of the existence of these two peripheral signaling streams has also contributed to the development of clinical tests. In patients, high-frequency vibrational stimuli to the skull or air-conducted sound can be used to induce ocular vestibular-evoked myogenic potentials (oVEMPs) mainly originating from the utricular macula or cervical VEMPs arising from the saccular macula [47]. These tests are applied in clinical practice to delineate otolith dysfunction in patients with posttraumatic dizziness, inferior vestibular neuritis or superior canal dehiscence syndrome [48, 49]. Experiments in rat and guinea pig have shown that this stimulation predominantly

activates irregular afferents, which are also responsive to bone-conducted or airborne sound [47]. Similarly, studies in monkey, chinchilla and rat have shown that galvanic vestibular stimulation (GVS) preferentially activates irregular afferents [39, 50]. Based on the non-invasiveness of the latter method and easy application in patients, GVS has found its entry into clinical practice, and facilitates a diagnosis and dissociation between Menierés disease and, e.g., vestibular migraine [51].

Vestibular neuronal typology as basis for sensory–motor processing

Central vestibular neurons play a key role in sensory–motor transformation of semicircular canal and otolith signals. Electrophysiological *in vitro* experiments in slices and isolated brain preparations have found two primary neuronal subtypes in the vestibular nucleus of rodents and guinea pig (type A and B neurons). These neurons distinctly differ in their resting discharge regularity, response dynamics and sets of ionic conductances [52]. While extensively characterized in guinea pig, rat and mouse, a similar dichotomy has been reported in frog (i.e., phasic and tonic neurons: [53]) and chicken (i.e., principle and elongate cells: [54]). Thus, a distinction into two vestibular subtypes appears to be the common denominator that matches the vestibular afferent organization [55].

Studies in behaving monkeys have established two main functional classes of vestibular neurons (Fig. 1a) that likely overlap with the classification scheme established *in vitro* [44]. The first class of neurons encodes voluntary eye movements as well as head motion. These Position-Vestibular-Pause (PVP) neurons project to extraocular motoneurons (Fig. 1a). A second class of neurons encodes head but not eye movements (i.e., vestibular-only (VO) neurons), projects to the spinal cord and is thought to mediate vestibulo-spinal reflexes for posture control (Fig. 1a). The response dynamics of the two neuronal classes in primates suggest that they receive input from two parallel sensory information streams (Fig. 1a–c). A first one, mediated by regular vestibular afferents, contains information about the stimulus' detailed time course (stimulus estimation). The second one, mediated by irregular vestibular afferents, transmits information about the occurrence of high-frequency stimulus features (feature detection). Notably, VO neurons show dynamic properties similar to irregular afferents and respond to the high-frequency features of motion stimuli in a strongly nonlinear fashion [56–58]. This behavior is similar to that of type B neurons that have been characterized *in vitro*. In contrast, PVP neurons show less dynamic properties, similar to regular afferents or to type A neurons described above. However, while various properties of type A and B neurons, obtained

in vitro, match very well with those of the two types of vestibular neurons recorded *in vivo*, to date a definitive functional description of both type A and B neurons during motion stimulation in the intact animal is lacking.

Structure for function: Similarities versus differences in central vestibular organization

Semicircular canal and otolith convergence

As a general vertebrate pattern, afferent projections from vestibular endorgans largely overlap in the different central vestibular nuclei of all investigated vertebrate species including cat [59], pigeon [60], frog [61] or fish [62, 63]. This demonstrates an evolutionarily inherent absence of a sensory map. At variance with the large overlap of afferent fibers from individual endorgans, however, second-order vestibular neurons (2°VN) are organized in a hindbrain segmental (rhombomeric) pattern that is based on the motor/premotor target of a particular vestibular subgroup [5, 64]. This rhombomeric arrangement was first demonstrated in chick embryos [65, 66], and further specified in fish [67], frog [68] and mouse [69]. The discovery of this segment-specific arrangement was established by visualization of Hox gene expression patterns as well as the plain visibility of rhombomeres in various vertebrate embryos or larvae that allowed the direct mapping of distinct vestibular subgroups onto the hindbrain scaffold.

Although afferents from all vestibular endorgans overlap to a large extent in the vestibular nuclei, 2°VNs exhibit a remarkable specificity in selecting their monosynaptic afferent input; most 2°VNs receive monosynaptic inputs from only one semicircular canal (Fig. 1d–f) or one otolith organ, respectively [21]. Moreover, semicircular canal inputs combine monosynaptically with otolith afferent inputs in a spatially specific manner. For example, horizontal semicircular canal signals predominantly converge with utricular signals (Fig. 1f) and vertical semicircular canal signals with those from vertical otolith organs [70]. Thus, despite the theoretical availability of sensory signals from all vestibular endorgans, the spatial motion vector of individual afferents is largely preserved at the level of the first neuronal element in the brainstem.

In everyday life, our vestibular sensors are activated by complex multi-dimensional motion patterns that simultaneously stimulate semicircular canal and otolith organs [26, 71]. Single-unit recordings in primates have further shown that integration of semicircular canal and otolith inputs by vestibular neurons is sub-additive and characterized by frequency-dependent (nonlinear) weighting of both modalities [72]. This integration is required to discriminate tilt from translation and to ensure accurate perception and motor performance.

Vestibular-visual convergence

The optokinetic reflex (OKR) works together with the vestibulo-ocular reflex (VOR) to stabilize gaze. The OKR is symmetric in monkeys and humans but asymmetric in mice, gerbils and rabbits [73–75] as well as in non-mammalian species such as frogs [21]. The symmetry in primates is likely mediated by relatively stronger contributions from cortical versus subcortical pathways [76–78]. Studies in behaving mammals indicate that eye-movement-sensitive vestibular nucleus neurons command OKR eye movements and also contribute to a “velocity storage” circuitry. This network uses visual information to supplement the decaying signal from vestibular afferents during sustained head movements to encode self-motion [79, 80]. Specifically, PVP but not VO neurons are modulated during both optokinetic and vestibular stimulation in mice [81] and potentially in primates [82].

Vestibular-proprioception convergence

Most vestibular nuclei neurons in mice [83], rats (e.g., [84]), cats [85–87] and alert squirrel and cynomolgus monkeys [88, 89] respond robustly to passive proprioceptive stimulation as well as to vestibular inputs. Neuronal responses to combined stimulation can generally be predicted based on the linear sum of a given neurons’ individual vestibular and proprioceptive sensitivities. However, there are important differences across species. While proprioceptive responses are robust in rodents, they are less pronounced in cynomolgus monkeys [89] and completely absent in rhesus monkeys [90, 91], a difference that is likely related to species-specific adaptations in gaze strategies during exploratory behavior.

Multimodal convergence during active self-motion

Recent neurophysiological studies in primates have emphasized the importance of extra-vestibular signals in shaping sensorimotor transformations that mediate vestibulo-spinal reflexes [44, 92]. Whereas the sensitivity or firing rate of vestibular nerve afferents is virtually identical during active or passive movements (Fig. 2a), the discharge of VO neurons shows striking differences in the two conditions. Notably, while VO neurons robustly respond during passive head movements, their vestibular-related modulation is markedly attenuated during active head movements (Fig. 2b). This attenuation is behaviorally advantageous during voluntary movements, since intact vestibulo-spinal reflexes would likely be counterproductive, eliciting postural responses that would oppose intended voluntary movements.

Neural circuits for the control of gaze: different constraints versus common principles

The circuitry underlying the VOR is the best characterized vestibular-driven pathway. In his classical studies, Lorente de Nó [93] demonstrated the existence of a three-neuronal arc that represents the most direct pathway between vestibular afferents and eye muscles. Thereafter, studies using many different vertebrate species to probe different aspects of this reflex circuitry, have established that the basic functional organization of the VOR has remained virtually unchanged since it evolved in early vertebrates [64]. In fact, this reflex pathway is remarkably well conserved in vertebrates spanning from fish to mammals with regard to segmentally arranged neuronal phenotypes, employed neurotransmitters, differential organization of the horizontal and vertical angular VOR and the conjugation of eye movements [5].

The relative simplicity of the VOR makes it an excellent model system for studies that try to bridge the gap between neuronal circuits and behavior. It is arguably our fastest behavior [94], consistent with the synaptic and axonal delays of the three-neuronal arc. The VOR shows a remarkable compensatory gain (eye velocity/head velocity = 1) as well as minimal phase lag over the physiological relevant range of head movements [94, 95]. The results of single-unit recordings in monkeys have provided insight into how the VOR effectively stabilizes gaze across a wide range of head velocities and frequencies experienced in everyday life [44]. Experiments in cats, monkeys and humans have shown that the efficacy of the VOR depends on the actual behavioral goal. While the VOR is compensatory when the goal is to stabilize gaze, it is attenuated or even suppressed when gaze (eye/head and or body movements) is redirected toward a target of interest.

The discovery of VOR efficacy as a function of the behavioral goal has particularly benefitted from single-unit analyses of multimodal integration in vestibular neurons of monkeys [44]. While vestibular afferents robustly encode head motion regardless of behavioral goals, responses of vestibular nucleus neurons (i.e., PVPs) are attenuated during gaze redirection (Fig. 2a). The neurophysiological bases of this suppressive influence are the well-characterized inhibitory projections from the brainstem premotor saccadic and pursuit pathways to the vestibular nuclei. Because an efference copy of the motor command to voluntarily redirect gaze suppresses the responses of PVP neurons, the efficacy of the VOR pathway critically depends on the actual gaze strategy during a particular situation (Fig. 2a). A more general role of motor efference copies in gaze stabilization is suggested by the findings in amphibians where copies of spinal locomotor commands

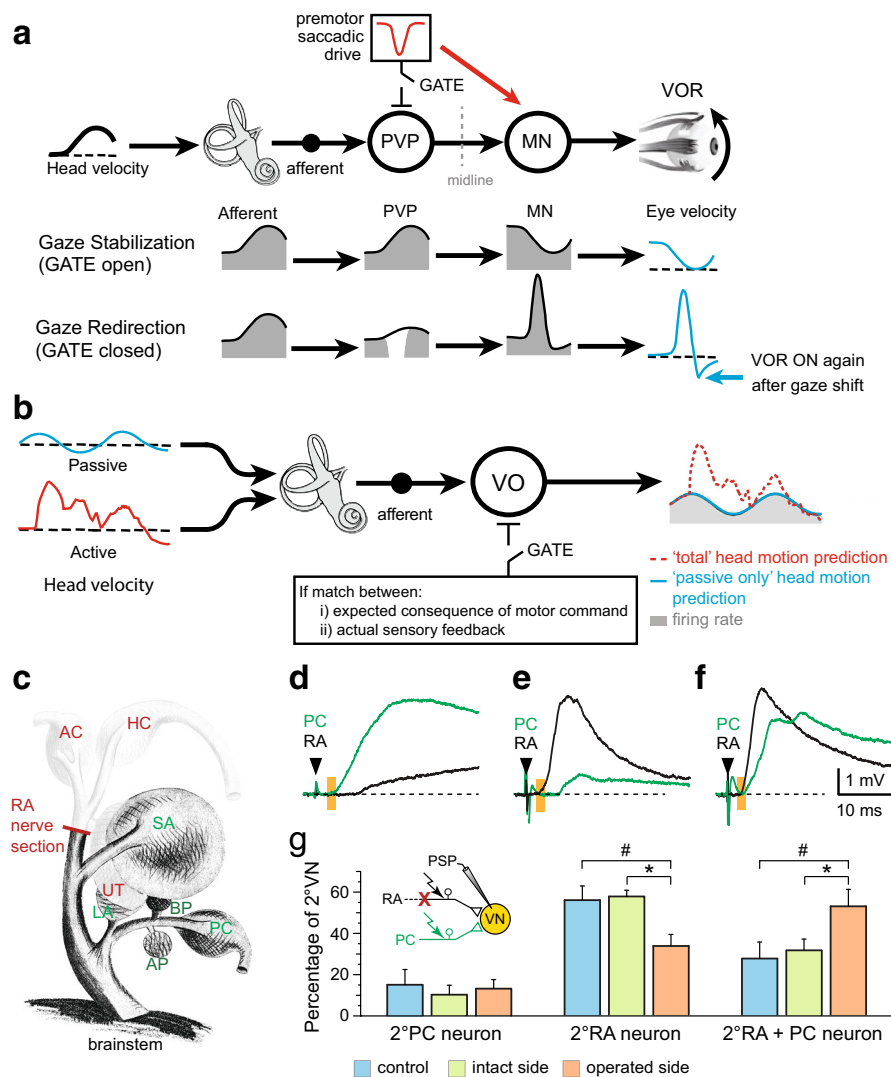


Fig. 2 Task-dependent and lesion-induced plasticity of vestibulo-ocular reflexes during active and passive motion. **a** Schematic illustrating task-specific processing (gaze stabilization, redirection) in central vestibular PVP neurons and extraocular motoneurons. **b** Task-specific cancellation of vestibular sensory inputs by predictive signals during active (red) head motion in VO neurons. In contrast, vestibular sensory information is completely transmitted during passive (blue) head motion. **c** Schematic view of the frog VIIIth nerve with endorgans, nerve branches and site of RA nerve section. **d–f** Convergence of afferent inputs from the RA and PC nerve branches; monosynaptic responses were evoked in some 2°VN after stimulation

of the PC nerve (green trace; **d**), in others after stimulation of the RA nerve (black trace; **e**) and in a third group after stimulation of both branches (**f**). **g** Percentages of the three types differ between controls and operated frogs and between intact and operated sides (color-coded bars). Black arrowhead and orange bars in **d–f** indicate stimulus and monosynaptic onset. AC, PC, HC anterior, posterior vertical, horizontal semicircular canal, AP, BP amphibian, basal papilla, LA lagena, RA ramus anterior of the VIIIth nerve, SA saccule, UT utricle, VN vestibular nuclei, VO vestibular-only neuron. **c–g** is based on data from [107]

directly drive compensatory eye movements during active motion [96, 97]. Given the neuronal connectivity between the spinal rhythm generator and the extraocular motor nuclei, it is very plausible that such connectivity represents an evolutionarily ancient pathway that might still have functional relevance in mammalian species. However, independent of its relevance in other vertebrates, it offers insight into a basic conceptual design that is important for understanding the role of the vestibular system in general.

In clinical practice, VOR testing has become highly important to detect deficits of the semicircular canals. The clinical head impulse test (HIT) for the horizontal semicircular canal was described for the first time by Halmagyi and Curthoys [98]. Today, video-based horizontal and vertical HIT recordings are routinely applied to measure VOR gain of all three semicircular canals to potentially detect compensatory saccades (so-called covert/overt saccades) in patients with vestibular disorders [99, 100]. VOR

testing thereby complements caloric irrigation, because both methods test different frequency spectra of vestibular afferents (high versus low frequency). In the clinical context, VOR testing is most important to differentiate a peripheral versus central origin of an acute vestibular syndrome [101, 102].

Principles of vestibular compensation and motor learning

Scientists and clinicians have long been fascinated by the capability of animals and humans to recover from the behavioral deficits after a loss of labyrinthine function [3, 10]. In the absence of a regrowth of the sensory organ, any functional improvement must be due to a reorganization of the signal processing in the central nervous system. Our current knowledge of the physiology of the compensation process has greatly benefitted from the employment of different animal models [103–105]. For example, work in rodents and guinea pigs has revealed a reduction of GABAergic commissural inhibition on the impaired side, changes in the strength of cerebellar inputs to the vestibular nuclei, as well as a shift toward more linear response properties of the deafferented type B vestibular neurons and an inverse pattern on the intact side [55]. Experiments in monkeys suggest a small but significant relative increase in the proportion of irregular afferents of the vestibular nerve on the contralesional side that parallels the shift in response properties observed in rodents and guinea pigs [106]. Experiments in frogs have further shown that a partial lesion of the vestibular nerve (Fig. 2c) causes synaptic reorganization of remaining ipsilateral afferent inputs onto deafferented 2°VNs (Fig. 2d–g) and a considerable reduction of the commissural inhibition at the expense of a modified vestibular reflex directional specificity [104, 107]. Across a variety of mammalian species including humans, monkeys, cats, guinea pigs and mice, VOR compensation is nearly complete for rotations toward the contralesional side, but incomplete for rotations toward the ipsilesional side, particularly for more dynamically challenging stimuli [105, 108]. Consistent with behavioral responses, experiments in behaving monkeys have shown that the responses of PVP neurons decrease immediately following a unilateral vestibular loss, but subsequently recover within a few weeks to reach values close to those measured before the lesion [109].

Findings across a variety of animal models have further established that a common denominator of vestibular compensation is the induction of homeostatic plasticity. For example, postural recovery in frogs after unilateral labyrinthectomy depends on an altered efficacy of spinal reflexes provided that body-weight-supporting limb proprioceptive inputs are available [55]. Interestingly, while this is the case in terrestrial vertebrates, aquatic species

lack these signals and instead develop scoliotic deformations likely due to a permanently manifested vestibular asymmetry [110]. Similarly, compensation in monkeys is mediated by rapid dynamic reweighting of inputs from different modalities (i.e., extra-vestibular proprioception and motor efference copy signals *versus* vestibular signals) at the level of vestibular nucleus neurons [109, 111–113]. Thus, multimodal integration is dynamically regulated in the vestibular system, in a manner that suggests a causal role for homeostatic plasticity in VOR compensation. This strategy appears common across vertebrates, providing a neural substrate for rehabilitation approaches currently used by clinicians to treat patients. Potential strategies for physical therapy after unilateral vestibular damage thus may include activities such as the Cawthorne-Cooksey exercises that involve a progression of increasingly complex head and body movements (reviewed in [114]), active VOR gain adaptation [115], neck muscle vibration [116] or an increased use of visual reference frames [117].

Plasticity within vestibular pathways also plays an essential role in fine-tuning the coordination and accuracy of the VOR in response to environmental or developmental alterations. For example, adaptive changes in VOR performance are required to compensate for the mismatch between visual and vestibular stimuli caused by the magnification of corrective lenses worn during common visual conditions [118, 119]. Such a gain control of the VOR is implemented by the prominent feed-forward cerebellar circuitry [21]. This circuit is highly plastic and provides the basis for the cerebellar contribution to motor learning. Experiments in cats and monkeys have further established that plasticity within the floccular complex of the cerebellum initially drives VOR adaptation, which in turn triggers longer term synaptic changes in floccular target neurons within the vestibular nuclei [120, 121]. In addition, *in vivo* and *in vitro* studies suggest that synaptic plasticity occurs within non-cerebellar VOR pathways alongside synaptic changes within the cerebellum [122–125]. Thus, it is likely that plasticity processes for gain modifications of vestibular reflexes are distributed features at multiple sites that allow guiding adaptations to maintain VOR accuracy.

Higher level processing and perception

Vestibular information is not only required for reflexive motor reactions but also vital for cognitive functions such as perception of self-motion, spatial orientation and body representation [126]. Single-unit studies in non-human primates have provided insight into the computations performed by the cerebellum and cortex. In addition, neuroimaging using caloric and galvanic vestibular stimulation have provided insight into how these higher order areas process vestibular stimuli [127].

The vestibular cerebellum integrates vestibular and extra-vestibular information to make fundamental contributions to self-motion perception. The nodulus–uvula (lobules X and IX) is thought to create an internal model of spatial orientation that accounts for the physics of our world. Consistent with theoretical predictions [128], some neurons combine otolith and semicircular canal inputs to distinguish tilt from translation [129]. In addition, the vestibular cerebellum integrates vestibular and proprioceptive inputs to represent head and body-in-space motion in two separate streams [130]. Specifically, neurons in the deep cerebellar nuclei (i.e., fastigial), which receive inputs from the anterior cerebellar vermis, encode body motion independently of head motion. Moreover, these neurons selectively and dynamically encode passive head and body motion relative to space, suggesting that the cerebellum computes an internal model of the expected sensory consequences of active head motion to selectively cancel respective responses [131]. This mechanism is likely responsible for the attenuation of active motion observed in early vestibular processing (i.e., see Fig. 2b).

Ascending projections from the vestibular nuclei and vestibular cerebellum terminate in regions of the thalamus (reviewed in [132, 133]), which in turn project to the cortex. In contrast to most other sensory systems, there is not a single primary cortical area devoted to vestibular signals. Instead, vestibular-related activity is found in multiple regions, including the parieto-temporal, frontal, somatosensory and extrastriate visual cortices (reviewed in [134]). Notably, most neurons in these areas receive converging visual and/or somatosensory inputs. Among these areas, the parietoinsular vestibular cortex (PIVC) is generally considered as primary vestibular cortex since (i) PIVC neurons respond to vestibular input [135–137], (ii) stimulation of this area produces vestibular sensations in humans [138], (iii) lesion of PIVC impairs perception of the subjective visual vertical [139] and (iv) cerebral blood flow of the PIVC area increases during vestibular stimulation [140–142]. Numerous studies have also focused on how vestibular processing in the dorsal medial superior temporal cortex (MSTd) contributes to our perception of self-motion (reviewed in [143]). The transmission of self-motion information from these cortical areas to entorhinal and perirhinal cortices and the hippocampus is thought to play a critical role in spatial cognition and navigation (reviewed in [144]). In fact, patients with a bilateral vestibulopathy show deficits in spatial orientation along with a markedly reduced hippocampal volume [145]. The contribution of the vestibular system in the pathophysiology of disorders of spatial attention such as neglect is increasingly recognized and can be used for rehabilitation [146, 147].

Advantages of different animal models

As illustrated in the previous chapters, studies using a wide variety of vertebrate species have been essential for furthering our knowledge of how the sensory organs in the inner ear detect head motion in space, how receptor cells transduce motion into voltage signals, and how the brain encodes and integrates these motion-related signals for accurate behavior and perception during self-motion. Notably, while some species are better suited than others to answer scientific questions on the organization and role of the vestibular system, others are better suited to establish the neural circuitry mediating higher order functions such as vestibular cognition. Vestibular research has particularly benefitted from its positioning at the intersection between basic and clinical science. For example, neural pathways mediating vestibular-driven reflex behaviors such as the VOR are relatively simple, and collective knowledge from studies using a wide variety of species have provided an excellent framework for understanding the physiology underlying clinical syndromes.

Overall, we argue that in the future a reasonably balanced spectrum of animal models will continue to be required to increase our understanding of the vestibular system. Studies of ontogenetic aspects and developmental assembly of appropriate neural connections will require further work in model species that allow easy access to embryonic stages. For instance, genetic approaches and developmental manipulations can be combined with physiology in vertebrates such as fish, frog, chicken or mouse, even though recordings are rather difficult to perform in embryonic and early post-embryonic birds or mammals [29]. With respect to cellular details of vestibulo-motor signal processing, electrophysiological recordings in slice preparations of rodents and guinea pigs have revealed important fundamental principles [55] and new innovative approaches including multichannel electrophysiological recordings and optical imaging will provide greater accessibility to population coding in vestibular structures. In addition, complementary studies in more “exotic” species will continue to contribute to reveal general concepts of sensory–motor transformation in vertebrates such as the Axolotl with its legendary regenerative capacity [148], turtles that allow robust *in vitro* experiments in an amniote vertebrate species [149], or flatfish that exhibit a substantial VOR reorganization during the transition from bilateral-symmetric free-swimming larvae to asymmetric bottom-dwelling adults [150].

For studies of evolution and adaptation in the vestibular system and its cellular components, as well as the respective computations performed by early vertebrate ancestors or higher order cortical processing, an even wider range of

vertebrate species is required [4, 67]. This is because central processing of vestibular signals depends on both intrinsic membrane and emerging network properties. Accordingly, deciphering their interactions requires experimental models with intact nervous systems that also provide the researcher with an experimental accessibility necessary to manipulate the respective neural circuitry. In the past, the isolated guinea pig or frog whole brain [55] substantially contributed to our understanding of underlying computations, with further improved probing of morpho-physiological aspects in recently developed semi-intact amphibian preparations [151]. We speculate that future mouse or zebrafish lines with genetically expressed calcium ion sensors will yield further enhanced accessibility to vestibulo-motor networks for in vivo recordings of cell and circuit activity.

In contrast, work in more advanced mammalian species including non-human primates will be required to further our understanding of how these circuits give rise to perception, cognition and behavior under normal conditions—knowledge essential for developing more effective health protocols to diagnose and treat the debilitating symptoms of vestibular disorders in patients. Decades of electrophysiological observations in non-human primates have already provided key insights as to how vestibular-driven sets of motor behavior and perception arise from neural circuit activity (reviewed in [44, 92]). Notably, primates will be the model of choice for studying the mechanisms that provide perceptual stability and accurate motor performance during common yet complex behaviors, such as combinations of voluntary head motion and locomotion. Further, while some basic brain circuits are preserved, many of the neural circuits related to higher cognitive functions differ between non-mammalian species and humans. By comparison, since the brain organization of humans and non-human primates are remarkably similar, especially with regard to the cerebellum and cerebral cortex, this animal model is particularly well suited for studies aimed at understanding the higher level organization of vestibular processing and motion perception [44, 92, 143, 152].

Finally, it is important to emphasize that studies across a wider range of species will be required to facilitate translational vestibular research progress. For studies focused on efficient, high-throughput drug discovery [153], understanding general circuit organization or multimodal interactions and the impact of motor efference copies for sensory–motor transformations [110], non-mammalian species are highly relevant and convenient model systems. In addition, drug treatments initially developed in species such as mice can fail when translated to humans, emphasizing the value of non-human primates in translational research. In summary, a comparative approach based on studies across a variety of vertebrate species, each with

particular advantages for defined scientific questions, remains necessary to maximize our understanding of the vestibular system and its pathophysiology.

Acknowledgments The authors thank M. Jamali and H. Dietrich for critically reading the manuscript. The authors acknowledge financial support by the German Federal Ministry of Education and Research under the Grant code 01 EO 0901 (to H.S. and A.Z.), Canadian Institutes of Health Research (to KEC) and National Institutes of Health Research (NIH) grant DC002390 (to K.E.C.). The authors declare that they have no conflict of interest.

Compliance with ethical standards

Conflicts of interest None.

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References

1. Churchland A, Lisberger S (2015) Contributions from different model organisms to brain research: introduction. *Neuroscience* 296:1–2
2. Goltz F (1870) Ueber die physiologische Bedeutung der Bogengänge des Ohrlabyrinths. *Pflügers Arch* 3:172–192
3. Ewald JR (1892) Physiologische Untersuchungen über das Endorgan des Nervus Octavus. JF Bergmann, Wiesbaden
4. Fritzsche B, Straka H (2014) Evolution of mechanosensory hair cells and inner ears: identifying stimuli to select altered molecular development toward new morphologies. *J Comp Physiol* 200:5–18
5. Straka H, Fritzsche B, Glover JC (2014) Connecting ears to eye muscles: evolution of a ‘simple’ reflex arc. *Brain Behav Evol* 83:162–175
6. Eugène D, Deforges S, Guimont F, Idoux E, Vidal PP, Moore LE, Vibert N (2007) Developmental regulation of the membrane properties of central vestibular neurons by sensory vestibular information in the mouse. *J Physiol* 583:923–943
7. Jones SM, Jones TA (2014) Genetics of peripheral vestibular dysfunction: lessons from mutant mouse strains. *J Am Acad Audiol* 25:289–301
8. Luebke AE, Holt JC, Jordan PM, Wong YS, Caldwell JS, Cullen KE (2014) Loss of α -calcitonin gene-related peptide (α CGRP) reduces the efficacy of the vestibulo-ocular reflex (VOR). *J Neurosci* 34:10453–10458
9. Flourens PP (1824) Recherches expérimentales sur les propriétés et les fonctions du système nerveux, dans les animaux vertébrés. Crevot, Paris
10. von Bechterew W (1883) Ergebnisse der Durchschneidung des N. acusticus nebst Erörterung der Bedeutung der semicirculären Kanäle für das Körpergleichgewicht. *Pflügers Arch*. 30:312–347
11. Crum-Brown A (1874) On the sense of rotation and the anatomy and physiology of the semicircular canals of the internal ear. *J Anat Physiol* 8:327–331
12. Breuer J (1874) Über die Funktion der Bogengänge des Ohrlabyrinths. *Wien Med Jahrb* 4:72–124

13. Mach E (1875) Grundlinien der Lehre von den Bewegungsempfindungen. W Engelmann, Leipzig
14. Webb WM (1905) The ear of fishes. Knowledge n ser 2:59–61
15. Parker GH (1903) The sense of hearing in fishes. Am Naturalist 37:185–204
16. Ach N (1901) Ueber die Otolithenfunction und den Labyrinthonus. Arch Ges Physiol 86:122–146
17. Parker GH (1908) Structure and functions of the ear of the squeteague. US Bur Fisheries Bull 28:1213–1224
18. Angelaki DE, Cullen KE (2008) Vestibular system: the many facets of a multimodal sense. Annu Rev Neurosci 31:125–150
19. Rabbitt RD, Damiano ER, Grant JW (2004) Biomechanics of the vestibular semicircular canals and otolith organs. In: Highstein SM, Fay RR, Popper AN (eds) The Vestibular System. Springer, New York, pp 153–201
20. Simpson JJ, Graf W (1985) The selection of reference frames by nature and its investigators. Rev Oculomot Res 1:3–16
21. Straka H, Dieringer N (2004) Basic organization principles of the VOR: lessons from frogs. Prog Neurobiol 73:259–309
22. Fritzsche B (1992) The water-to-land transition: evolution of the tetrapod basilar papilla, middle ear, and auditory nuclei. In: Webster DB, Fay RR, Popper AN (eds) The Evolutional Biology of Hearing. Springer, New York, pp 351–375
23. Ladham A, Pickles JO (1996) Morphology of the monotreme organ of Corti and macula lagena. J Comp Neurol 366:335–347
24. Wu LQ, Dickman JD (2011) Magnetoreception in an avian brain in part mediated by inner ear lagena. Curr Biol 21:418–423
25. Jones GM, Spells KE (1963) A theoretical and comparative study of the functional dependence of the semicircular canal upon its physical dimensions. Proc R Soc Lond B Biol Sci 157:403–419
26. Schneider AD, Jamali M, Carriot J, Chacron MJ, Cullen KE (2015) The increased sensitivity of irregular peripheral canal and otolith vestibular afferents optimizes their encoding of natural stimuli. J Neurosci 35:5522–5536
27. Eatock RA, Songer JE (2011) Vestibular hair cells and afferents: two channels for head motion signals. Annu Rev Neurosci 34:501–534
28. Steinacker A (2004) Sensory Processing and Ionic Currents in Vestibular Hair Cells. In: Highstein SM, Fay RR, Popper AN (eds) Springer handbook of auditory research. Springer, New York, pp 202–234
29. Straka H (2010) Ontogenetic rules and constraints of vestibulo-ocular reflex development. Curr Opin Neurobiol 20:689–695
30. Muller M (1999) Size limitations in semicircular duct systems. J Theor Biol 198:405–437
31. Rabbitt RD (1999) Directional coding of three-dimensional movements by the vestibular semicircular canals. Biol Cybern 80:417–431
32. Hullar TE (2006) Semicircular canal geometry, afferent sensitivity, and animal behavior. Anat Rec A Discov Mol Cell Evol Biol 288:466–472
33. Yang A, Hullar TE (2007) Relationship of semicircular canal size to vestibular-nerve afferent sensitivity in mammals. J Neurophysiol 98:3197–3205
34. Lambert FM, Beck JC, Baker R, Straka H (2008) Semicircular canal size determines the developmental onset of angular vestibuloocular reflexes in larval *Xenopus*. J Neurosci 28:8086–8096
35. Beck JC, Gilland E, Tank DW, Baker R (2004) Quantifying the ontogeny of optokinetic and vestibuloocular behaviors in zebrafish, medaka, and goldfish. J Neurophysiol 92:3546–3561
36. Branoner F, Straka H (2014) Semicircular canal-dependent developmental tuning of translational vestibulo-ocular reflexes in *Xenopus laevis*. Dev Neurobiol. doi:10.1002/dneu.22234
37. Baird RA, Lewis ER (1986) Correspondences between afferent innervation patterns and response dynamics in the bullfrog utricle and lagena. Brain Res 369:48–64
38. Lewis ER, Li CW (1975) Hair cell types and distributions in the otolithic and auditory organs of the bullfrog. Brain Res 83:35–50
39. Goldberg JM (2000) Afferent diversity and the organization of central vestibular pathways. Exp Brain Res 130:277–297
40. Baird RA (1994) Comparative transduction mechanisms of hair cells in the bullfrog utricle. II. Sensitivity and response dynamics to hair bundle displacement. J Neurophysiol 71:685–705
41. Vranceanu F, Perkins GA, Terada M, Chidavaenzi RL, Ellisman MH, Lysakowski A (2012) Striated organelle, a cytoskeletal structure positioned to modulate hair-cell transduction. Proc Natl Acad Sci USA 109:4473–4478
42. Holstein GR, Rabbitt RD, Martinelli GP, Friedrich VL Jr, Boyle RD, Highstein SM (2004) Convergence of excitatory and inhibitory hair cell transmitters shapes vestibular afferent responses. Proc Natl Acad Sci USA 101:15766–15771
43. Hullar TE, Della Santina CC, Hirvonen T, Lasker DM, Carey JP, Minor LB (2005) Responses of irregularly discharging chinchilla semicircular canal vestibular-nerve afferents during high-frequency head rotations. J Neurophysiol 93:2777–2786
44. Cullen KE (2012) The vestibular system: multimodal integration and encoding of self-motion for motor control. Trends Neurosci 35:185–196
45. Jamali M, Carriot J, Chacron MJ, Cullen KE (2013) Strong correlations between sensitivity and variability give rise to constant discrimination thresholds across the otolith afferent population. J Neurosci 33:11302–11313
46. Sadeghi SG, Chacron MJ, Taylor MC, Cullen KE (2007) Neural variability, detection thresholds, and information transmission in the vestibular system. J Neurosci 27:771–781
47. Curthoys IS (2010) A critical review of the neurophysiological evidence underlying clinical vestibular testing using sound, vibration and galvanic stimuli. Clin Neurophysiol 121:132–144
48. Kim JS, Kim HJ (2012) Inferior vestibular neuritis. J Neurol 259:1553–1560
49. Brantberg K, Verrecchia L (2009) Testing vestibular-evoked myogenic potentials with 90-dB clicks is effective in the diagnosis of superior canal dehiscence syndrome. Audiol Neurotol 14:54–58
50. Goldberg JM, Smith CE, Fernández C (1984) Relation between discharge regularity and responses to externally applied galvanic currents in vestibular nerve afferents of the squirrel monkey. J Neurophysiol 51:1236–1256
51. Neuhauser HK (2007) Epidemiology of vertigo. Curr Opin Neurol 20:40–46
52. Vidal PP, Vibert N, Serafin M, Babalian A, Mühlethaler M, de Waele C (1999) Intrinsic physiological and pharmacological properties of central vestibular neurons. Adv Otorhinolaryngol 55:26–81
53. Straka H, Beranek M, Rohregger M, Moore LE, Vidal PP, Vibert N (2004) Second-order vestibular neurons form separate populations with different membrane and discharge properties. J Neurophysiol 92:845–861
54. Peusner KD, Giaume C (1997) Ontogeny of electrophysiological properties and dendritic pattern in second-order chick vestibular neurons. J Comp Neurol 384:621–633
55. Straka H, Vibert N, Vidal PP, Moore LE, Dutia MB (2005) Intrinsic properties of vertebrate vestibular neurons: function, development and plasticity. Prog Neurobiol 76:349–392
56. Goldberg JM, Highstein SM, Moschovakis AK, Fernández C (1987) Inputs from regularly and irregularly discharging vestibular nerve afferents to secondary neurons in the vestibular

- nuclei of the squirrel monkey. I. An electrophysiological analysis. *J Neurophysiol* 58:700–718
57. Highstein SM, Goldberg JM, Moschovakis AK, Fernández C (1987) Inputs from regularly and irregularly discharging vestibular nerve afferents to secondary neurons in the vestibular nuclei of the squirrel monkey. II. Correlation with output pathways of secondary neurons. *J Neurophysiol* 58:719–738
 58. Massot C, Schneider AD, Chacron MJ, Cullen KE (2012) The vestibular system implements a linear-nonlinear transformation in order to encode self-motion. *PLoS Biol* 10:e1001365
 59. Gacek RR (1969) The course and central termination of first order neurons supplying vestibular endorgans in the cat. *Acta Otolaryngol Suppl* 254:1–66
 60. Dickman DJ, Fang Q (1996) Differential central projections of vestibular afferents in pigeons. *J Comp Neurol* 367:110–131
 61. Birinyi A, Straka H, Matesz C, Dieringer N (2001) Location of dye-coupled second order and of efferent vestibular neurons labeled from individual semicircular canal or otolith organs in the frog. *Brain Res* 921:44–59
 62. McCormick CA, Braford MR Jr (1994) Organization of inner ear endorgan projections in the goldfish, *Carassius auratus*. *Brain Behav Evol* 43:189–205
 63. McCormick CA, Wallace AC (2012) Otolith end organ projections to auditory neurons in the descending octaval nucleus of the goldfish, *Carassius auratus*: a confocal analysis. *Brain Behav Evol* 80:41–63
 64. Straka H, Baker R (2013) Vestibular blueprint in early vertebrates. *Front Neural Circuits* 7:182
 65. Glover JC, Petursdottir G (1991) Regional specificity of developing reticulospinal, vestibulospinal and vestibulo-ocular projections in the chicken embryo. *J Neurobiol* 22:353–376
 66. Cambronerio F, Puelles L (2000) Rostrocaudal nuclear relationships in the avian medulla oblongata: a fate map with quail chick chimeras. *J Comp Neurol* 427:522–545
 67. Baker R (1998) From genes to behavior in the vestibular system. *Otolaryngol Head Neck Surg* 119:263–275
 68. Straka H, Baker R, Gilland E (2001) Rhombomeric organization of vestibular pathways in larval frogs. *J Comp Neurol* 437:42–55
 69. Pasqualetti M, Diaz C, Renaud JS, Rijli F, Glover JC (2007) Fate-mapping the mammalian hindbrain: segmental origins of vestibular projection neurons assessed using rhombomere-specific *hoxa2* enhancer elements in the mouse embryo. *J Neurosci* 27:9670–9681
 70. Straka H, Holler S, Goto F (2002) Patterns of canal and otolith afferent input convergence in frog second order vestibular neurons. *J Neurophysiol* 88:2287–2301
 71. Carriot J, Brooks JX, Cullen KE (2013) Multimodal integration of self-motion cues in the vestibular system: active versus passive translations. *J Neurosci* 33:19555–19566
 72. Carriot J, Jamali M, Brooks JX, Cullen KE (2015) Integration of canal and otolith inputs by central vestibular neurons is subadditive for both active and passive self-motion: implication for perception. *J Neurosci* 35:3555–3565
 73. Stahl JS, James RA, Oommen BS, Hoebeek FE, De Zeeuw CI (2005) Eye movements of the murine P/Q calcium channel mutant tottering, and the impact of aging. *J Neurophysiol* 95:1588–1607
 74. Kaufman GD (2002) Video-oculography in the gerbil. *Brain Res* 958:472–487
 75. Collewijn H (1981) Asymmetry of monocular optokinetic nystagmus (OKN) in the rabbit is not abolished by unilateral enucleation at birth. *Neurosci Lett Suppl* 7:461
 76. Schor CM (1983) Subcortical binocular suppression affects the development of latent and optokinetic nystagmus. *Am J Optom Physiol Opt* 60:481–502
 77. Atkinson J, Braddick O, Braddick F (1974) Acuity and contrast sensitivity of infant vision. *Nature* 247:403–404
 78. Zee DS, Tusa RJ, Herdman SJ, Butler PH, Gucer G (1987) Effects of occipital lobectomy upon eye movements in primate. *J Neurophysiol* 58:883–907
 79. Cohen B, Suzuki JJ, Raphan T (1983) Role of the otolith organs in generation of horizontal nystagmus: effects of selective labyrinthine lesions. *Brain Res* 276:159–164
 80. Wearne S, Raphan T, Cohen B (1998) Control of spatial orientation of the angular vestibuloocular reflex by the nodulus and uvula. *J Neurophysiol* 79:2690–2715
 81. Beraneck M, Cullen KE (2007) Activity of vestibular nuclei neurons during vestibular and optokinetic stimulation in the alert mouse. *J Neurophysiol* 98:1549–1565
 82. Bryan AS, Angelaki DE (2009) Optokinetic and vestibular responsiveness in the macaque rostral vestibular and fastigial nuclei. *J Neurophysiol* 101:714–720
 83. Medrea I, Cullen KE (2013) Multisensory integration in early vestibular processing in mice: the encoding of passive versus active motion. *J Neurophysiol* 110:2704–2717
 84. Barresi M, Grasso C, Li Volsi G, Manzoni D (2013) Effects of body to head rotation on the labyrinthine responses of rat vestibular neurons. *Neuroscience* 244:134–146
 85. Anastasopoulos D (1982) Mergner T Canal-neck interaction in vestibular nuclear neurons of the cat. *Exp Brain Res* 46:269–280
 86. Boyle R, Pompeiano O (1981) Convergence and interaction of neck and macular vestibular inputs on vestibulospinal neurons. *J Neurophysiol* 45:852–868
 87. Wilson VJ, Yamagata Y, Yates BJ, Schor RH, Nonaka S (1990) Response of vestibular neurons to head rotations in vertical planes. III. Response of vestibulocollic neurons to vestibular and neck stimulation. *J Neurophysiol* 64:1695–1703
 88. Gdowski GT, Belton T, McCrear RA (2001) The neurophysiological substrate for the cervico-ocular reflex in the squirrel monkey. *Exp Brain Res* 140:253–264
 89. Sadeghi SG, Mitchell DE, Cullen KE (2009) Different neural strategies for multimodal integration: comparison of two macaque monkey species. *Exp Brain Res* 195:45–57
 90. Roy JE, Cullen KE (2001) Selective processing of vestibular reafference during self-generated head motion. *J Neurosci* 21:2131–2142
 91. Roy JE, Cullen KE (2004) Dissociating self-generated from passively applied head motion: neural mechanisms in the vestibular nuclei. *J Neurosci* 24:2102–2111
 92. Cullen KE (2011) The neural encoding of self-motion. *Curr Opin Neurobiol* 21:587–595
 93. Lorente de Nó R (1933) Vestibulo-ocular reflex arc. *Arch Neurol Psychiatr (chic)* 30:245–291
 94. Huterer M, Cullen KE (2002) Vestibuloocular reflex dynamics during high-frequency and high-acceleration rotations of the head on body in rhesus monkey. *J Neurophysiol* 88:13–28
 95. Armand M, Minor LB (2001) Relationship between time- and frequency-domain analyses of angular head movements in the squirrel monkey. *J Comput Neurosci* 11:217–239
 96. Lambert FM, Combes D, Simmers J, Straka H (2012) Gaze stabilization by efference copy signaling without sensory feedback during vertebrate locomotion. *Curr Biol* 22:1649–1658
 97. von Uckermann G, Le Ray D, Combes D, Straka H, Simmers J (2013) Spinal efference copy signaling and gaze stabilization during locomotion in juvenile *Xenopus* frogs. *J Neurosci* 33:4253–4264
 98. Halmagyi GM, Curthoys IS (1988) A clinical sign of canal paresis. *Arch Neurol* 45:737–739
 99. Bartl K, Lehen N, Kohlbecher S, Schneider E (2009) Head impulse testing using video-oculography. *Ann N Y Acad Sci* 1164:331–333

100. MacDougall HG, Weber KP, McGarvie LA, Halmagyi GM, Curthoys IS (2009) The video head impulse test: diagnostic accuracy in peripheral vestibulopathy. *Neurology* 73:1134–1141
101. Cnyrim CD, Newman-Toker D, Karch C, Brandt T, Strupp M (2008) Bedside differentiation of vestibular neuritis from central “vestibular pseudoneuritis”. *J Neurol Neurosurg Psychiatry* 79:458–460
102. Kattah JC, Talkad AV, Wang DZ, Hsieh YH, Newman-Toker DE (2009) HINTS to diagnose stroke in the acute vestibular syndrome: three-step bedside oculomotor examination more sensitive than early MRI diffusion-weighted imaging. *Stroke* 40:3504–3510
103. Curthoys IS (2000) Vestibular compensation and substitution. *Curr Opin Neurol* 13:27–30
104. Dieringer N (2003) Activity-related postlesional vestibular reorganization. *Ann NY Acad Sci* 1004:50–60
105. Cullen KE, Minor LB, Beraneck M, Sadeghi SG (2009) Neural substrates underlying vestibular compensation: contribution of peripheral versus central processing. *J Vestib Res* 19:171–182
106. Sadeghi SG, Minor LB, Cullen KE (2006) Response of vestibular-nerve afferents to active and passive rotations under normal conditions and after unilateral labyrinthectomy. *J Neurophysiol* 97:1503–1514
107. Goto F, Straka H, Dieringer N (2000) Expansion of afferent vestibular signals after the section of one of the vestibular nerve branches. *J Neurophysiol* 84:581–584
108. Smith PF, Curthoys IS (1989) Mechanisms of recovery following unilateral labyrinthectomy: a review. *Brain Res Brain Res Rev* 14:155–180
109. Sadeghi SG, Minor LB, Cullen KE (2010) Neural correlates of motor learning in the vestibulo-ocular reflex: dynamic regulation of multimodal integration in the macaque vestibular system. *J Neurosci* 30:10158–10168
110. Lambert FM, Straka H (2012) The frog vestibular system as a model for lesion-induced plasticity: basic neural principles and implications for posture control. *Front Neurol* 3:42
111. Sadeghi SG, Minor LB, Cullen KE (2011) Multimodal integration after unilateral labyrinthine lesion: single vestibular nuclei neuron responses and implications for postural compensation. *J Neurophysiol* 105:661–673
112. Sadeghi SG, Minor LB, V KE (2012) Neural correlates of sensory substitution in vestibular pathways following complete vestibular loss. *J Neurosci* 32:14685–14695
113. Jamali M, Mitchell DE, Dale A, Carrier J, Sadeghi SG, Cullen KE (2014) Neuronal detection thresholds during vestibular compensation: contributions of response variability and sensory substitution. *J Physiol* 592:1565–1580
114. Ricci NA, Aratani MC, Doná F, Macedo C, Caovilla HH, Ganança FF (2010) A systematic review about the effects of the vestibular rehabilitation in middle-age and older adults. *Rev Bras Fisioter* 14:361–371
115. Migliaccio AA, Schubert MC (2013) Unilateral adaptation of the human angular vestibulo-ocular reflex. *J Assoc Res Otolaryngol* 14:29–36
116. Strupp M, Arbusow V, Dieterich M, Sautier W, Brandt T (1998) Perceptual and oculomotor effects of neck muscle vibration in vestibular neuritis. Ipsilateral somatosensory substitution of vestibular function. *Brain* 121:677–685
117. Horak FB (2010) Postural compensation for vestibular loss and implications for rehabilitation. *Restor Neurol Neurosci* 28:57–68
118. Boyden ES, Katoh A, Raymond JL (2004) Cerebellum-dependent learning: the role of multiple plasticity mechanisms. *Annu Rev Neurosci* 27:581–609
119. Lisberger SG (2009) Internal models of eye movement in the floccular complex of the monkey cerebellum. *Neuroscience* 162:763–776
120. Cullen KE (2008) Procedural learning: VOR. In: Eichenbaum H (ed) *Memory Systems. Learning and Memory: A Comprehensive Reference*, Vol. 3, Academic Press/Elsevier, Oxford, pp 383–402
121. Broussard DM, Titley HK, Antflick J, Hampson DR (2011) Motor learning in the VOR: the cerebellar component. *Exp Brain Res* 210:451–463
122. McElvain LE, Bagnall MW, Sakatos A, du Lac S (2010) Bidirectional plasticity gated by hyperpolarization controls the gain of postsynaptic firing responses at central vestibular nerve synapses. *Neuron* 68:763–775
123. Scarduzio M, Panichi R, Pettorossi VE, Grassi S (2012) The repetition timing of high frequency afferent stimulation drives the bidirectional plasticity at central synapses in the rat medial vestibular nuclei. *Neuroscience* 223:1–11
124. Beraneck M, McKee JL, Aleisa M, Cullen KE (2008) Asymmetric recovery in cerebellar-deficient mice following unilateral labyrinthectomy. *J Neurophysiol* 100:945–958
125. Zwergal A, Schlichtiger J, Xiong G, Beck R, Günther L, Schniepp R, Schöberl F, Jahn K, Brandt T, Strupp M, Bartenstein P, Dieterich M, Dutia MB, la Fougère C (2014) Sequential [18F]FDG μ PET whole-brain imaging of central vestibular compensation: a model of deafferentation-induced brain plasticity. *Brain Struct Funct*. doi:10.1007/s00429-014-0899-1
126. Mast FW, Preuss N, Hartmann M, Grabherr L (2014) Spatial cognition, body representation and affective processes: the role of vestibular information beyond ocular reflexes and control of posture. *Front Integr Neurosci* 8:44. doi:10.3389/fnint.2014.00044
127. Dieterich M, Brandt T (2008) Functional brain imaging of peripheral and central vestibular disorders. *Brain* 131:2538–2552
128. Merfeld D, Zupan L, Peterka R (1999) Humans use internal models to estimate gravity and linear acceleration. *Nature* 398:615–618
129. Laurens J, Meng H, Angelaki DE (2013) Neural representation of orientation relative to gravity in the macaque cerebellum. *Neuron* 80:1508–1518
130. Brooks JX, Cullen KE (2009) Multimodal integration in rostral fastigial nucleus provides an estimate of body movement. *J Neurosci* 29:10499–10511
131. Brooks JX, Cullen KE (2013) The primate cerebellum selectively encodes unexpected self-motion. *Curr Biol* 23:947–955
132. Lopez C, Blanke O (2011) The thalamocortical vestibular system in animals and humans. *Brain Res Rev* 67:119–146
133. Kirsch V, Keiser D, Hergenroeder T, Erat O, Ertl-Wagner B, Brandt T, Dieterich M (2015) Structural and functional connectivity mapping of the vestibular circuitry from human brainstem to cortex. *Brain Struct Funct*. doi:10.1007/s00429-014-0971-x
134. Guldin WO, Grüsser OJ (1998) Is there a vestibular cortex? *Trends Neurosci* 21:254–259
135. Grüsser OJ, Pause M, Schreier U (1990) Vestibular neurones in the parieto-insular cortex of monkeys (*Macaca fascicularis*): visual and neck receptor responses. *J Physiol* 430:559–583
136. Chen et al (2014) Chen X, DeAngelis GC, Angelaki DE (2014) Eye-centered visual receptive fields in the ventral intraparietal area. *J Neurophysiol* 112:353–361
137. Shinder ME, Newlands SD (2014) Sensory convergence in the parieto-insular vestibular cortex. *J Neurophysiol* 111:2445–2464
138. Penfield W (1957) Vestibular sensation and the cerebral cortex. *Ann Otol Rhinol Laryngol*. 66:691–698

139. Brandt T, Dieterich M, Danek A (1994) Vestibular cortex lesions affect the perception of verticality. *Ann Neurol* 35:403–412
140. Friberg L, Olsen TS, Roland PE, Paulson OB, Lassen NA (1985) Focal increase of blood flow in the cerebral cortex of man during vestibular stimulation. *Brain* 108:609–623
141. Stephan T, Deutschländer A, Nolte A, Schneider E, Wiesmann M, Brandt T, Dieterich M (2005) Functional MRI of galvanic vestibular stimulation with alternating currents at different frequencies. *Neuroimage* 26:721–732
142. Schlindwein P, Mueller M, Bauermann T, Brandt T, Stoeter P, Dieterich M (2008) Cortical representation of saccular vestibular stimulation: vEMPs in fMRI. *Neuroimage* 39:19–31
143. Angelaki DE, Gu Y, De Angelis GC (2011) Visual and vestibular cue integration for heading perception in extrastriate visual cortex. *J Physiol* 589:825–833
144. Hitier M, Besnard S, Smith PF (2014) Vestibular pathways involved in cognition. *Front Integr Neurosci* 8:59
145. Brandt T, Schautzer F, Hamilton DA, Brüning R, Markowitsch HJ, Kalla R, Darlington C, Smith P, Strupp M (2005) Vestibular loss causes hippocampal atrophy and impaired spatial memory in humans. *Brain* 128:2732–2741
146. Brandt T, Dieterich M, Strupp M, Glasauer S (2012) Model approach to neurological variants of visuo-spatial neglect. *Biol Cybern* 106:681–690
147. Wilkinson D, Zubko O, Sakel M, Coulton S, Higgins T, Pullicino P (2014) Galvanic vestibular stimulation in hemi-spatial neglect. *Front Integr Neurosci* 8:4. doi:[10.3389/fnint.2014.00004](https://doi.org/10.3389/fnint.2014.00004)
148. Dörenberger S, Banchi R, Brosel S, Seebacher C, Laimgruber S, Uhl R, Felmy F, Straka H, Kunz L (2015) Analysis of signal processing in vestibular circuits with a novel light-emitting diodes-based fluorescence microscope. *Eur J Neurosci* 41:1332–1344
149. Highstein SM, Mann MA, Holstein GR, Rabbitt RD (2015) The quantal component of synaptic transmission from sensory hair cells to the vestibular calyx. *J Neurophysiol* 113:3827–3835
150. Graf W, Baker R (1983) Adaptive changes of the vestibulo-ocular reflex in flatfish are achieved by reorganization of central nervous pathways. *Science* 221:777–779
151. Straka H, Simmers J (2012) *Xenopus laevis*: an ideal experimental model for studying the developmental dynamics of neural assembly and sensory motor computations. *Dev Neurobiol* 72:649–663
152. Fetsch CR, DeAngelis GC, Angelaki DE (2013) Bridging the gap between theories of sensory cue integration and the physiology of multisensory neurons. *Nat Rev Neurosci* 14:429–442
153. Wheeler GN, Brändli AW (2009) Simple models for chemical genetics and drug discovery screens: lessons from zebrafish and *Xenopus*. *Dev Dyn* 238:1287–1308